

# Half-cell Study of La and Ca doped Strontium Titanates Anode for Direct Methane Solid Oxide Fuel Cell

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## Abstract

Solid oxide fuel cell (SOFC) has been providing high conversion efficiency of chemical energy to electricity without any pollution. One of the major advantages of SOFC over other fuel cell is use of direct natural gas at high temperature without any external reformer. Conventional nickel-yttria stabilized zirconia (Ni-YSZ) composite anode provides excellent catalytic property, current collection and stability for H<sub>2</sub> oxidation but it is not tolerant towards sulphur poisoning and also accelerates coke deposition in presence of hydrocarbon fuels. It necessitates the use of alternate anode for direct hydrocarbon fuel. In the present work, attempts have been made to apply La and Ca doped A-site deficient SrTiO<sub>3</sub> (LSCT<sub>A-</sub>) as potential anode for direct methane SOFC. Low catalytic activity of LSCT<sub>A-</sub> is improved by infiltration of Ni and CeO<sub>2</sub> catalyst. Half cell (YSZ/4%Ni-6%CeO<sub>2</sub>-LSCT<sub>A-</sub>) provided 200 mW cm<sup>-2</sup> maximum power density and regain its initial performance in H<sub>2</sub> even after 6 h exposure to humidified CH<sub>4</sub> at 800 °C.

## Introduction

Solid oxide fuel cell (SOFC) is a promising power device in scenario of current global environmental issues and increased demand for alternative energy sources. SOFC is highly efficient and require no external reformer as it directly converts hydrocarbon fuel to electrical energy. Oxidation of fuel takes place at anode so choice of high catalytic active and efficient anode material is the most important consideration in direct hydrocarbon SOFC. Conventionally, nickel (Ni) is widely used as anode because of its excellent catalytic property, current collection and stability towards H<sub>2</sub> oxidation but it is not tolerant to sulphur poisoning and also accelerates coke deposition in presence of hydrocarbon fuels. Coke deposition drastically degrades the SOFC performance. Significant reduction in the coke deposition using anodes such as Cu-CeO<sub>2</sub> and Fe-CeO<sub>2</sub> has been observed (1-3). Thermal instability and limited electrochemical activity of copper based anodes at SOFC operation temperature need use of alternate anode materials for sustained stable performance in presence of direct hydrocarbon fuels (4-5). Agglomeration of Ni particles due to sintering leads to cell deactivation during operation at high temperature (800 °C), which is another major

concern. Transition Metal oxides such as  $\text{TiO}_2$ ,  $\text{CeO}_2$  and  $\text{Nb}_2\text{O}_5$  in small amount (~5-10 wt %) are added to Ni-YSZ anode to prevent performance degradation due to Ni coarsening (6-9). Recently, single and double doped  $\text{SrTiO}_3$  (both at A and B site) has been researched extensively and it exhibited that doped  $\text{SrTiO}_3$  not only prevent the loss due Ni sintering but also provides a excellent alternate to Ni based anode for hydrocarbon fuels. Higher valent cation such as  $\text{La}^{3+}$  when substituted at  $\text{Sr}^{2+}$  site induce semiconducting behavior in  $\text{SrTiO}_3$ . The substitution of  $\text{Sr}^{2+}$  with  $\text{La}^{3+}$  can be compensated by introducing the extra oxygen beyond the  $\text{ABO}_3$  stoichiometry under oxidizing conditions and maintains the electro-neutrality. Strontium titanate ( $\text{SrTiO}_3$ ) possesses desirable thermal and chemical stability and show mixed ionic and electronic conducting (MIEC) behavior at low oxygen partial pressure (10-12). It is shown that A-site deficiency enhances the electrical conductivity of  $\text{La}^{3+}$  doped  $\text{SrTiO}_3$  (13-15). Ca doping lowers down the sintering temperature of  $\text{LST}_A$ - backbone and enhances the reducibility and electrical conductivity (13). Catalytic activity of  $\text{LSCT}_A$ - towards fuel oxidation has been improved by infiltration of transition metal and metal oxides such as Cu, Ni and  $\text{CeO}_2$  (13, 16-17).

In the current work, A-site deficient  $\text{La}_{0.2}\text{Sr}_{0.25}\text{Ca}_{0.45}\text{TiO}_3$  ( $\text{LSCT}_A$ -) anode backbone is fabricated, characterized and used in electrolyte supported half-cells.  $\text{LSCT}_A$ - backbone is impregnated with nitrate precursors of Ni and/or  $\text{CeO}_2$  to increase the catalytic activity and the cell performances in  $\text{H}_2$  and humidified  $\text{CH}_4$  at 800 °C (18-19). The present work explores the potential of La and Ca co-doped A-site deficient  $\text{SrTiO}_3$  anode for direct methane SOFC.

## Experimental

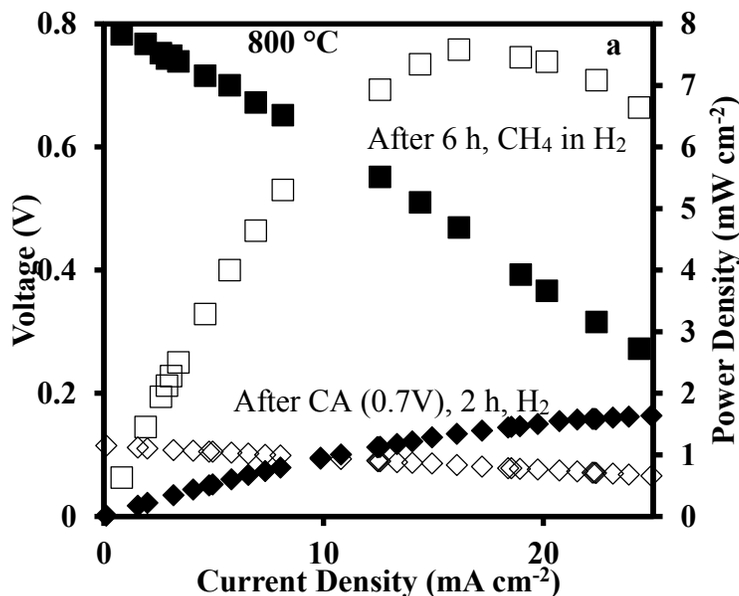
Aqueous tape casting and co-sintering of dense YSZ electrolyte and porous  $\text{La}_{0.2}\text{Sr}_{0.25}\text{Ca}_{0.45}\text{TiO}_3$  ( $\text{LSCT}_A$ -) anode has been carried out to fabricate half-cells (18-19). The  $\text{LSCT}_A$ - powder for anode was mixed with graphite (pore former) and milled for 24 h by adding de-ionized water as solvent, hypermer KD 6 as dispersant. Poly(ethylene glycol), glycerol as plasticizer, poly vinyl alcohol and ethoxylated 2,4,7,9-tetra ethyl 5 decyn-4,7-diol as deformer were added to the slurry and milled for next 24 h. After that, suspension was de-aired and cast to get green tapes. Similarly, dense YSZ green tapes were casted. Green tapes of  $\text{LSCT}_A$ - and YSZ were laminated together to get dense YSZ/porous  $\text{LSCT}_A$ -. Then green laminated tapes were cut in to appropriate size and co-sintered at 1350 °C. The sintered anode was reduced in  $\text{H}_2$  atmosphere at 950 °C.  $\text{CeO}_2$  and/or Ni catalyst were impregnated in  $\text{LSCT}_A$ - porous anode matrix using  $\text{Ce}(\text{NO}_3)_3 \cdot 6\text{H}_2\text{O}$  and  $\text{Ni}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$  solution (18-19). Multiple impregnation and calcination at 500 °C were carried out till required loading in porous  $\text{LSCT}_A$ - was achieved. Silver wire current collectors were applied to both the electrodes using silver paste and active area of cell was 0.5  $\text{cm}^2$ . Cell was sealed onto ceramic tube using Aremco, Ceramabond 552. Hydrogen ( $\text{H}_2$ ) or humidified methane ( $\text{CH}_4$ ) gas was supplied with flow rate of 50 mL/min to the anode. The current-voltage characteristics followed by impedance studies at open circuit voltage (OCV) were carried out using potentiostat/galvanostat (Autolab, Metrohm). The microstructure, morphology and elemental mapping of half-cells before and after cell operation had been analyzed using scanning electron microscope (SEM) and energy dispersive x-ray spectroscopy (EDX) using Zeiss EVO 50. The

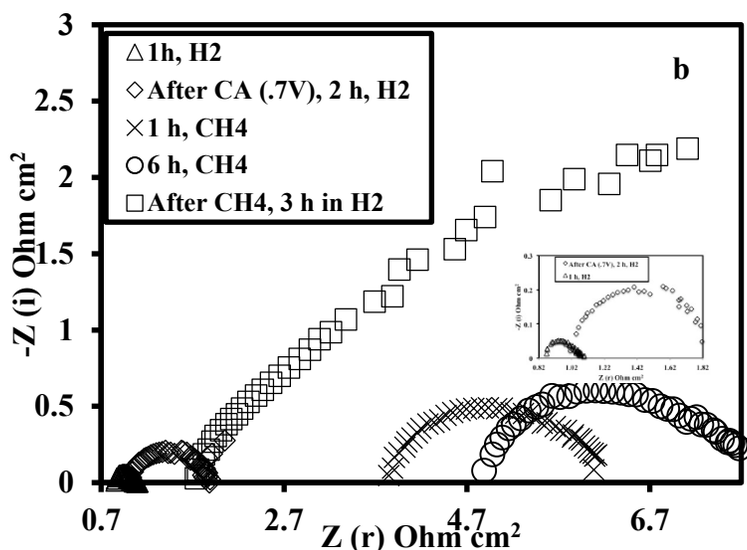
particle size and selected area diffraction patterns were observed using HR-TEM (TEM, Technai G<sup>2</sup> 200 KV, FEI).

## Result and Discussion

### Electrolyte supported half-cell testing

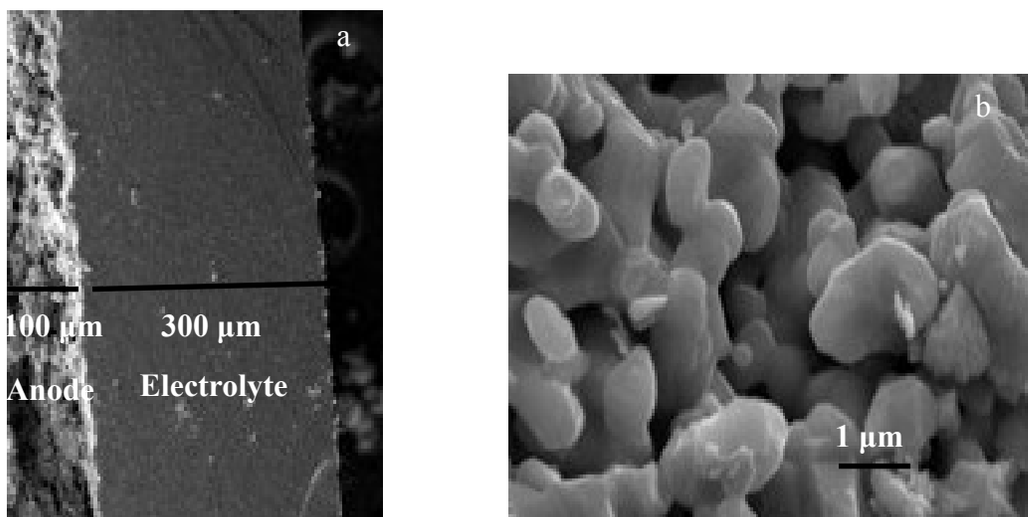
Electrolyte (YSZ) supported half-cell with LSCT<sub>A</sub>- as anode has been electrochemically characterized in H<sub>2</sub> and humidified CH<sub>4</sub> fuel at 800 °C to assess the electro catalytic activity of LSCT<sub>A</sub>- (fig. 1a-b). Cell is operated in H<sub>2</sub> fuel at 800 °C for 2 h and very low open circuit voltage (OCV) is observed. To activate the anode, chronoamperometry (CA) at 0.7 V has been carried out for 45 minute and OCV increases up to 0.1 V, which is still very low. After 3 h in H<sub>2</sub>, humidified CH<sub>4</sub> is supplied as fuel to anode chamber. The OCV of cell slowly increases and after 6 h exposure to CH<sub>4</sub> in presence of H<sub>2</sub> fuel, 0.8 V OCV is recorded. LSCT<sub>A</sub>- anode shows poor catalytic activity towards H<sub>2</sub> and CH<sub>4</sub> oxidation as cell performance in terms of current and power density is very low as presented in fig. 1a. At 1 h of cell operation in H<sub>2</sub>, 0.86 ohm cm<sup>2</sup> ohmic resistance is observed which increases to 4.89 ohm cm<sup>2</sup> at 6 h of cell operation in humidified CH<sub>4</sub> but decreases down to 1.735 ohm cm<sup>2</sup> in presence of H<sub>2</sub> fuel after 6 h exposure of CH<sub>4</sub>. The polarisation resistance increases from 0.212 at 1 h in H<sub>2</sub> to 2.60 ohm cm<sup>2</sup> at 6 h operation in CH<sub>4</sub>. Inset of fig. 1b shows the impedance spectra corresponds to 1 h operation in H<sub>2</sub> and after CA (0.7V) at 2 h of operation. It can be infer from above mentioned analyses that LSCT<sub>A</sub>- is catalytically inactive for methane decomposition but it does not get deteriorated due to the coke deposition during operation as evident from the decrement in resistance and enhanced OCV in H<sub>2</sub> atmosphere after CH<sub>4</sub> exposure for 6 h. Carbon particles provides the conducting pathway instead of blocking the active surface hence better result in H<sub>2</sub> after CH<sub>4</sub> exposure. To increase the catalytic activity of anode backbone, impregnation of the porous LSCT<sub>A</sub>- with active metal catalyst has been carried out and corresponding results are presented here.



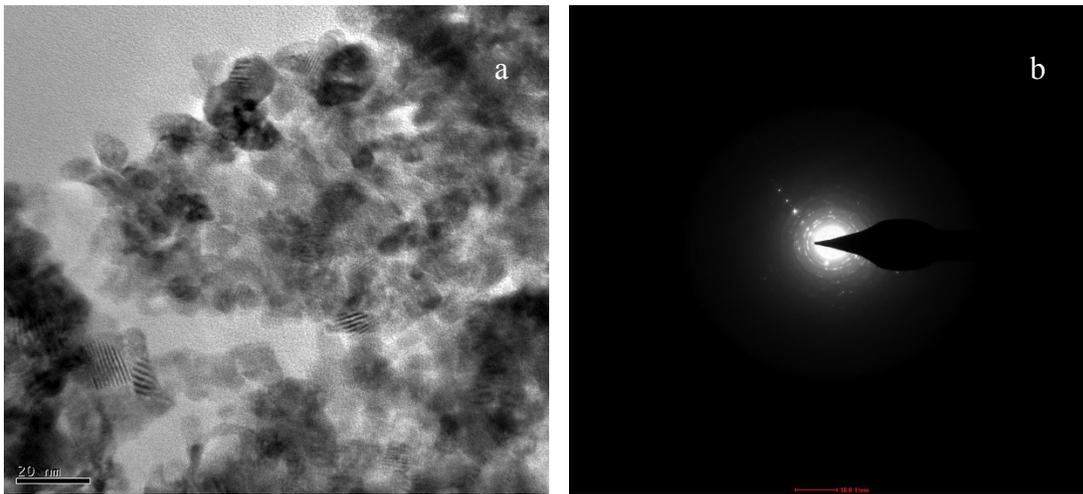


**Fig. 1 (a) i-V characteristics, (b) Impedance spectra at OCV of half cell (YSZ/LSCT<sub>A</sub>-) in H<sub>2</sub> and humidified CH<sub>4</sub> at 800 °C.**

Fig. 2(a-b) shows SEM of fracture surface and LSCT<sub>A</sub>- anode after cell testing. The thickness of electrolyte and anode are 300 and 100 μm respectively. Electrolyte is perfectly dense and proper adherence between electrolyte and anode is observed even after 12 h of cell operation at 800 °C. Fig 2b shows well connected grains of LSCT<sub>A</sub>- with sufficient porosity in stable anode backbone structure. Average particle size is ~ 1-2 μm. After cell testing HRTEM of LSCT<sub>A</sub>- is carried out. Fig. 3 shows HRTEM of agglomerated LSCT<sub>A</sub>- particles. Diffraction patterns correspond to crystalline nature of LSCT<sub>A</sub>- is shown in the fig. 3a. SAED pattern in fig 3b shows regular crystal diffraction pattern corresponds to LSCT<sub>A</sub>- crystal.



**Fig. 2 (a) SEM of (a) Fracture surface and (b) Anode of half cell (YSZ/LSCT<sub>A</sub>-) after testing in H<sub>2</sub> and humidified CH<sub>4</sub>.**



**Fig. 3 (a) HRTEM and (b) SAED of LSCT<sub>A</sub>- after testing in H<sub>2</sub> and humidified CH<sub>4</sub>.**

To enhance the catalytic activity, 6% CeO<sub>2</sub> is impregnated in porous LSCT<sub>A</sub>- anode and tested in electrolyte supported half-cell in H<sub>2</sub> and humidified CH<sub>4</sub> at 800 °C. Fig. 4(a-b) presents the i-V, i-P and impedance spectra of YSZ/6%CeO<sub>2</sub>-LSCT<sub>A</sub>- half cell. As shown in fig. 4a, maximum power density 132 mW cm<sup>-2</sup> (305 mA cm<sup>-2</sup>) is observed at 1 h in H<sub>2</sub> fuel. After 3 h of cell operation in H<sub>2</sub>, humidified CH<sub>4</sub> is fed as fuel. Maximum power density in CH<sub>4</sub> after 6 h is 70 mW cm<sup>-2</sup> (174 mA cm<sup>-2</sup>), which is lower than that of the power density observed in H<sub>2</sub>. The half cell regains its initial performance once the H<sub>2</sub> is again fed and maximum power density 111 mW cm<sup>-2</sup> (305 mA cm<sup>-2</sup>) has been observed. The performance degradation in CH<sub>4</sub> is due to sluggish reaction associated with the dissociation of CH<sub>4</sub>. It is significant to note that degradation in performance due to CH<sub>4</sub> is not drastic and reasonable performance is shown by cell even after 6 h exposure of CH<sub>4</sub> during operation at 800 °C. It has been observed from impedance analyses, shown in fig. 4b that ohmic as well as polarisation resistance in H<sub>2</sub> is not changed significantly before and after 6 h operation in humidified CH<sub>4</sub>. Ohmic and total polarization in H<sub>2</sub> and CH<sub>4</sub> changed from 0.832 to 1.531 ohm cm<sup>2</sup> and 0.38 to 0.7574 ohm cm<sup>2</sup> respectively. In CH<sub>4</sub>, the ohmic as well as polarization resistance increased significantly which might be due to sluggish reaction kinetics corresponds to CH<sub>4</sub> oxidation on LSCT<sub>A</sub>- backbone. Presence of CeO<sub>2</sub> enhances the ionic conductivity as well as catalytic activity in anode backbone and does not accelerate the coke formation as infer from the above discussion. Morphological and elemental analysis of cell after 12 h of operation is presented in fig. 5(a-c). Well-distributed fine nano sized CeO<sub>2</sub> particles on LSCT<sub>A</sub>- backbone as well as in the pores can be seen from fig. 5a. HRTEM in fig. 5b shows the diffraction patterns correspond to LSCT<sub>A</sub>- crystal. SAED pattern shown in inset of fig. 5c confirms the crystalline CeO<sub>2</sub> along with the regular pattern corresponds to LSCT<sub>A</sub>-. Presence of constituent elements La, Sr, Ca, Ti, O and Ce in anode is confirmed from the EDX analysis after operation as presented in fig. 5c.

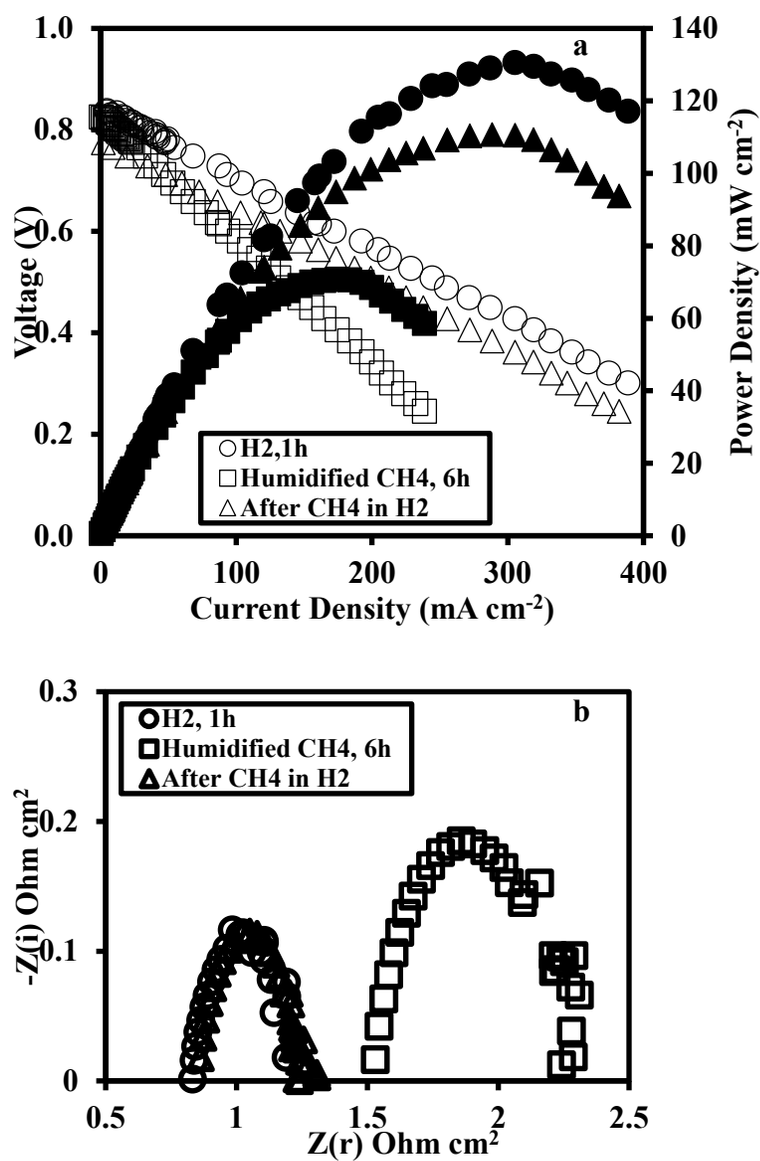
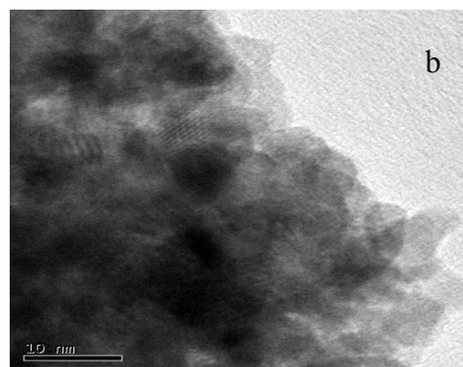
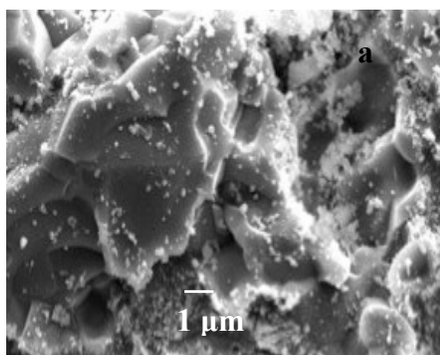
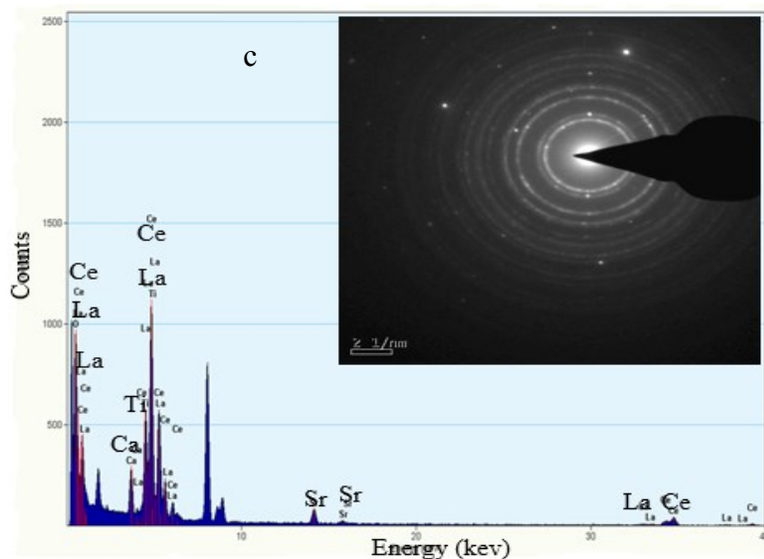


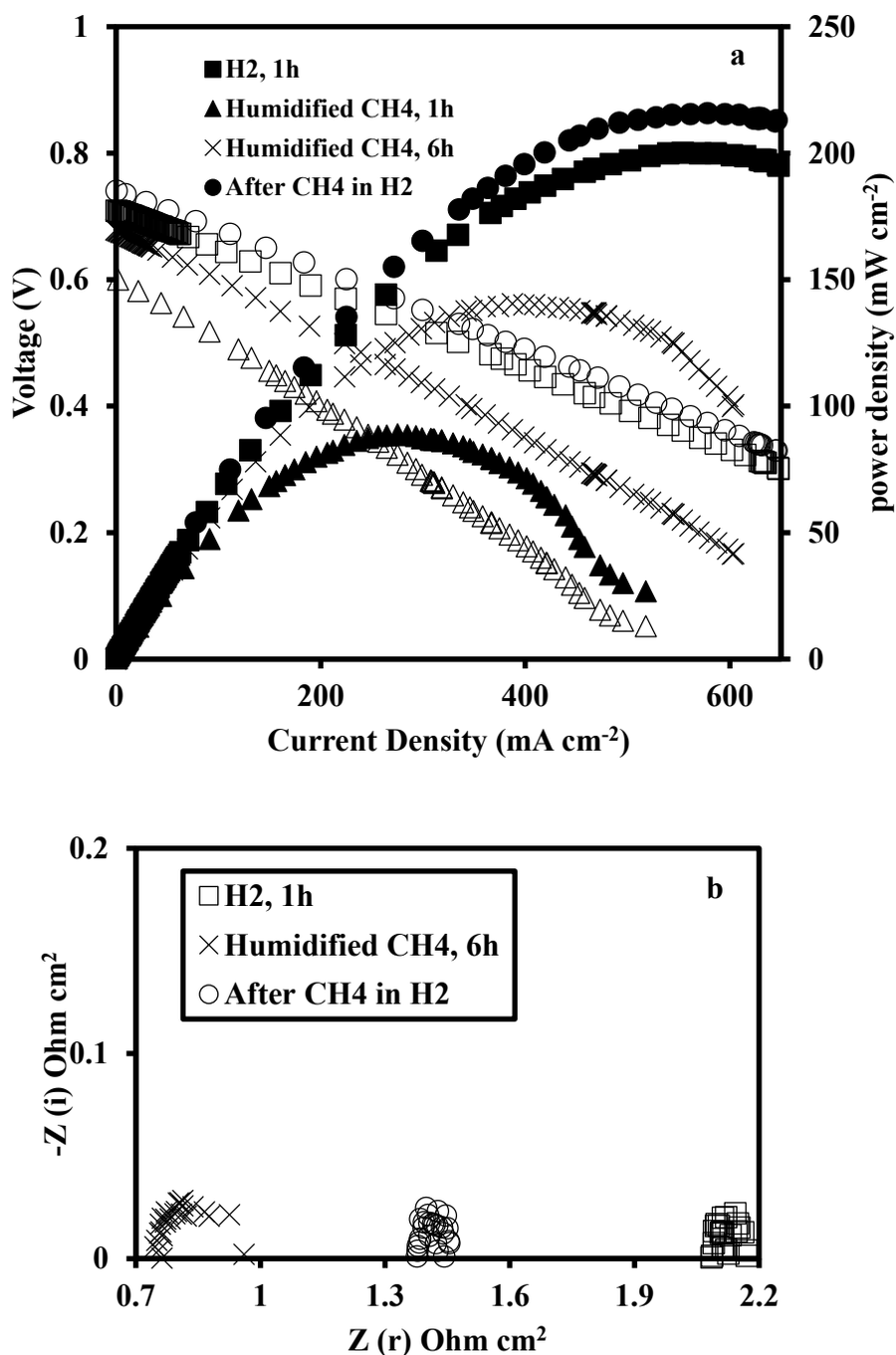
Fig. 4 (a) i-V characteristics, (b) Impedance spectra at OCV of half cell (YSZ/6%CeO<sub>2</sub>-LSCT<sub>A</sub>) in H<sub>2</sub> and humidified CH<sub>4</sub>, 800 °C.





**Fig. 5 (a) SEM, (b) HRTEM and, (c) Elemental analysis, SAED of anode of half cell (YSZ/6%CeO<sub>2</sub>-LSCT<sub>A</sub>) after testing in H<sub>2</sub> and humidified CH<sub>4</sub>.**

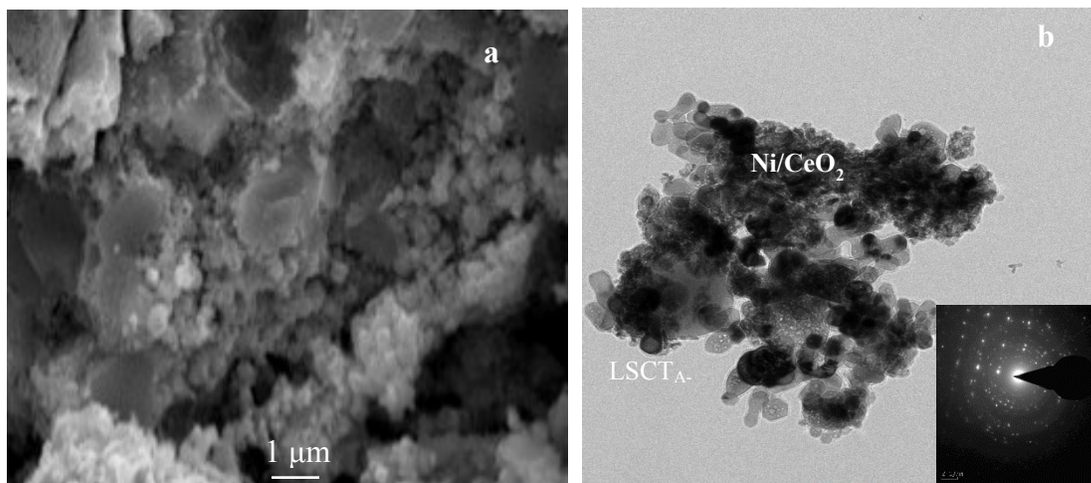
4% Ni is impregnated along with the 6% CeO<sub>2</sub> in porous LSCT<sub>A</sub> anode to enhance the half-cell performance by increasing catalytic activity. i-V, i-P and impedance analysis of half-cell shown in fig. 6 (a-b) explains the performance in H<sub>2</sub> as well as in CH<sub>4</sub> at 800 °C. Maximum power density of cell is 200 mW cm<sup>-2</sup> (554 mA cm<sup>-2</sup>) at 1 h of operation in H<sub>2</sub>, which increases up to 216 mW cm<sup>-2</sup> (578 mA cm<sup>-2</sup>) in H<sub>2</sub> after 6h of cell operation in humidied CH<sub>4</sub>. In humidified CH<sub>4</sub> exposure, maximum power density of cell is 89 mW cm<sup>-2</sup> (285 mA cm<sup>-2</sup>) at 1 h of operation, which increases up to 140 mW cm<sup>-2</sup> (409 mA cm<sup>-2</sup>) at 6 h of operation. At elevated temperature the LSCT<sub>A</sub> backbone along with the catalysts is getting reduced further hence the increase in performance with time has been observed. Improved performance also indicates that Ni particles in presence of CeO<sub>2</sub> are not get agglomerated due to sintering at elevated temperature. This phenomenon suggests strong metal support interaction (SMSI) between Ni and CeO<sub>2</sub>. Similar behavior between Ni and TiO<sub>2</sub> on YSZ backbone has been reported earlier (6-7). Lower performance in CH<sub>4</sub> is due to the sluggish reaction corresponding to CH<sub>4</sub> oxidation. So, it can be infer here that Ni is not promoting the coke deposition in presence of CeO<sub>2</sub>, as reasonable performance is observed in humidified CH<sub>4</sub> environment and even in H<sub>2</sub> atmosphere after CH<sub>4</sub> exposure for 6 h at 800 °C. i-V and i-P analyses are corroborated with the impedance results presented in fig. 6 b. It is observed that the ohmic as well polarization resistance decreases in H<sub>2</sub> before and after CH<sub>4</sub> exposure. Ohmic resistance decreases from 2.083 to 1.376 ohm cm<sup>2</sup> before and after 6 h operation in CH<sub>4</sub> respectively. The decrease in ohmic resistance may be due to deposition of carbon particles in anode matrix during operation in CH<sub>4</sub>.



**Fig. 6 (a) i-V characteristics, (b) Impedance spectra at OCV of half cell (YSZ/4% Ni-6% CeO<sub>2</sub>-LSCT<sub>A</sub>-) in H<sub>2</sub> and humidified CH<sub>4</sub>, 800 °C.**

Recently published work by researchers from Irvine group (19) reported enhanced performance of LSCT<sub>A</sub>- in presence of small amount of Ni and CeO<sub>2</sub> in H<sub>2</sub> atmosphere but they did not explore the viability of LSCT<sub>A</sub>- in CH<sub>4</sub>. Fig. 7 (a-b) represents the morphology, HRTEM and SAED pattern of cell after operation for 12 h. SEM micrograph (fig.7a) shows the well-dispersed Ni/CeO<sub>2</sub> anode catalyst on and in the pores of LSCT<sub>A</sub>-. Well-patterned grains of LSCT<sub>A</sub>- as well as finely dispersed Ni/CeO<sub>2</sub> particles on and around LSCT<sub>A</sub>- can be observed from fig.7b. It has been

observed that other than regular crystal diffraction patterns some separate patterns correspond to Ni/CeO<sub>2</sub> are also present in SAED as shown in the inset of fig. 7b.



**Fig. 7 (a) SEM, (b) HRTEM and SAED of anode of half cell (YSZ/4%Ni-6%CeO<sub>2</sub>-LSCT<sub>A</sub>-) after testing in H<sub>2</sub> and humidified CH<sub>4</sub>, 800 °C**

### Conclusion

Electrolyte supported half-cells with LSCT<sub>A</sub>- anode backbone have been tested in H<sub>2</sub> and humidified CH<sub>4</sub> fuel. Low catalytic activity of LSCT<sub>A</sub>- towards electro oxidation of fuel is significantly improves in presence of metallic catalyst. 6% CeO<sub>2</sub> loading in LSCT<sub>A</sub>- backbone improves the performance of the cell in H<sub>2</sub> as well as in humidified CH<sub>4</sub> at 800 °C. It has been observed that coke deposition did not degrade the anode active surface even after 6 h of exposure of CH<sub>4</sub> at elevated temperature. 4% Ni infiltration in 6% CeO<sub>2</sub>-LSCT<sub>A</sub>- enhances the performance from 131 mW cm<sup>-2</sup> to 200 mW cm<sup>-2</sup> at 800 °C in H<sub>2</sub>. In humidified CH<sub>4</sub> environment, cell having 4% Ni-6% CeO<sub>2</sub>-LSCT<sub>A</sub>- anode shows low performance (140 mW cm<sup>-2</sup> Power density) due to sluggish methane oxidation reaction but provided enhanced power density (216 mW cm<sup>-2</sup>) once H<sub>2</sub> is again fed after 6 h cell operation in humidified CH<sub>4</sub>. Performance of the cell is reasonably good as results presented here correspond to cells having thick electrolyte (300 μm). Cells are structurally robust even after 12 h of accelerated testing. LSCT<sub>A</sub>- is a potential anode backbone material for direct methane SOFC and could be an excellent anode for other hydrocarbons.

### References

1. R. J. Gorte and J. M. Vohs, *J. Catalysis*, **216**, 477 (2003).
2. S. K. Lee, A. Kipyung, J. M. Vohs and R. J. Gorte, *Electrochem. Solid-State Lett.*, **8**, A48 (2005).
3. G. Kaur and S. Basu, *J. Power Sources*, **241**, 783 (2013).
4. G. Kaur and S. Basu, *Fuel Cells*, **14**(6), 1006 (2014).

5. G. Kaur and S. Basu, *Int. J. Energy Res.*, **39**, 1345 (2015).
6. C. A. Singh, L. Bansal, P. Tiwari and V. V. Krishnan, *ECS Trans.*, **25**(2), 897 (2009).
7. P. Tiwari and S. Basu, *Int. J. Hydrogen Energy*, **38**, 9494 (2013).
8. P. Tiwari and S. Basu, *J. Solid State Electrochem.*, **18**, 805 (2014).
9. P. Tiwari and S. Basu, *ECS Trans.*, **57**(1), 1545 (2013).
10. C. Sun and U. Stimming, *J. Power Sources*, **171**, 247 (2007).
11. S. Tao and J. T. S. Irvine, *J. Mater. Chem.*, **12**, 2356 (2002).
12. S.W.Tao and J.T.S.Irvine, *Chemical Record*, **4**(2), 83 (2004).
13. A. D. Aljaberi and J.T.S. Irvine, *J. Mater. Chem. A*, **1**, 5868 (2013).
14. C. D. Savaniu and J.T.S. Irvine, *Solid State Ionics*, **192**, 491 (2011).
15. P. Tiwari and S. Basu, *ECS Trans.*, **68** (1), 1435 (2015).
16. A. Yaqub, C. D. Savaniu, N. K. Janjua and J. T. S Irvine, *J. Mater. Chem. A*, **1**, 14189 (2013).
17. M. C. Verbraeken, B. Iwanschitz, A. Mai and J. T. S. Irvine, *J. Electrochem. Soc.*, **159** (11), F757 (2012).
18. L. Lu, M. C. Verbraeken, M. Cassidy and J. T. S. Irvine, *ECS Trans.*, **57**(1), 1415 (2013).
19. L. Lu, C. Ni, M. Cassidy and J.T.S. Irvine, *J. Mater. Chem.*, **4**, 11708 (2016).